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A2

(54) Title: PROCESS FOR THE PREPARATION OF L-AMINO ACIDS USING STRAINS OF THE ENTEROBACTERIACEAE FAMILY

(57) Abstract: The invention relates to a process for the preparation of L-amino acids, in particular L-threonine, in which the following steps are carried out: a) fermentation of microorganisms of the Enterobacteriaceae family which produce the desired L-amino acid, b) separation of the culture medium from the microorganisms, c) extraction of the L-amino acid from the culture medium, d) purification of the L-amino acid by means of a column chromatography, e) crystallization of the L-amino acid, f) isolation of the L-amino acid, g) drying of the L-amino acid, h) storage of the L-amino acid, i) packaging of the L-amino acid, j) labeling of the L-amino acid, k) distribution of the L-amino acid, l) sale of the L-amino acid, m) delivery of the L-amino acid to the customer.

**Process for the Preparation of L-Amino Acids Using
Strains of the Enterobacteriaceae Family**

Field of the Invention

This invention relates to a process for the preparation of
5 L-amino acids, in particular L-threonine, using strains of
the Enterobacteriaceae family in which at least one or more
of the genes of the cysteine biosynthesis pathway (cysteine
biosynthetic pathway) chosen from the group consisting of
cysG, cysB, cysZ, cysK, cysM, cysA, cysW, cysU, cysP, cysD,
10 cysN, cysC, cysJ, cysI, cysH, cysE and sbp is (are)
enhanced.

Prior Art

L-Amino acids, in particular L-threonine, are used in human
medicine and in the pharmaceuticals industry, in the
15 foodstuffs industry and very particularly in animal
nutrition.

It is known to prepare L-amino acids by fermentation of
strains of Enterobacteriaceae, in particular Escherichia
coli (E. coli) and Serratia marcescens. Because of their
20 great importance, work is constantly being undertaken to
improve the préparation processes. Improvements to the
process can relate to fermentation measures, such as e.g.
stirring and supply of oxygen, or the composition of the
nutrient media, such as e.g. the sugar concentration during
25 the fermentation, or the working up to the product form, by
e.g. ion exchange chromatography, or the intrinsic output
properties of the microorganism itself.

Methods of mutagenesis, selection and mutant selection are
used to improve the output properties of these
30 microorganisms. Strains which are resistant to
antimetabolites, such as e.g. the threonine analogue α-
amino-β-hydroxyvaleric acid (AHV), or are auxotrophic for
arginine, are also mentioned, and it is also mentioned

acids, such as e.g. L-threonine, are obtained in this manner.

Methods of the recombinant DNA technique have also been employed for some years for improving the strain of 5 strains of the Enterobacteriaceae family which produce L-amino acids, by amplifying individual amino acid biosynthesis genes and investigating the effect on the production.

Object of the Invention

10 The object of the invention is to provide new measures for improved fermentative preparation of L-amino acids, in particular L-threonine.

Summary of the Invention

The invention provides a process for the fermentative 15 preparation of L-amino acids, in particular L-threonine, using microorganisms of the Enterobacteriaceae family which in particular already produce L-amino acids and in which at least one or more of the nucleotide sequence(s) which code(s) for the genes of the cysteine biosynthesis pathway 20 (cysteine biosynthetic pathway) chosen from the group consisting of cysG, cysB, cysZ, cysK, cysM, cysA, cysW, cysU, cysP, cysD, cysN, cysC, cysJ, cysI, cysH, cysE and sbp is (are) enhanced.

The process according to the invention for the preparation 25 of amino acids comprises the following steps:

- a) fermentation of the microorganisms of the Enterobacteriaceae family which produce the desired L-amino acid and in which one or more of the genes of the cysteine biosynthesis pathway chosen from the group consisting of cysG, cysB, cysZ, cysK, cysM, cysA, cysW, cysU, cysP, cysD, cysN, cysC, cysJ, cysI, cysH, cysE and sbp, or nucleotide

sequences which code for them is/are enhanced, in particular over-expressed,

- b) concentration of the desired L-amino acid in the medium or in the cells of the microorganisms, and
- 5 c) isolation of the desired L-amino acid, constituents of the fermentation broth and/or the biomass in its entirety or portions (> 0 to 100%) thereof optionally remaining in the product.

Detailed Description of the Invention

10 The use of endogenous genes is preferred. "Endogenous genes" or "endogenous nucleotide sequences" are understood as meaning the genes or nucleotide sequences present in the population of a species.

Where L-amino acids or amino acids are mentioned in the
15 following, this means one or more amino acids, including their salts, chosen from the group consisting of L-asparagine, L-threonine, L-serine, L-glutamate, L-glycine, L-alanine, L-cysteine, L-valine, L-methionine, L-isoleucine, L-leucine, L-tyrosine, L-phenylalanine, L-
20 histidine, L-lysine, L-tryptophan and L-arginine. L-Threonine is particularly preferred.

The term "enhancement" in this connection describes the increase in the intracellular activity or concentration of one or more enzymes or proteins in a microorganism which
25 are coded by the corresponding DNA, for example by increasing the number of copies of the gene or genes, using a potent promoter or a gene or allele which codes for a corresponding enzyme or protein with a high activity, and optionally combining these measures.

30 By enhancement measures, in particular over-expression, the activity or concentration of the corresponding protein is increased in a defined way at least 10%, 25%, 50%, 75% or 100%.

150%, 200%, 300%, 400% or 500%, up to a maximum of 1000% or 2000%, based on that of the wild-type protein or the activity or concentration of the protein in the starting microorganism.

5 The process comprises carrying out the following steps:

- a) fermentation of microorganisms of the Enterobacteriaceae family in which one or more of the genes of the cysteine biosynthesis pathway chosen from the group consisting of cysG, cysB, cysZ, cysK, cysM, cysA, cysW, cysU, cysP, cysD, cysN, cysC, cysJ, cysI, cysH, cysE and sbp is (are) enhanced,
- b) concentration of the corresponding L-amino acid in the medium or in the cells of the microorganisms of the Enterobacteriaceae family, and
- c) isolation of the desired L-amino acid, constituents of the fermentation broth and/or the biomass in its entirety or portions (> 0 to 100 %) thereof optionally remaining in the product.

20 The microorganisms which the present invention provides can produce L-amino acids from glucose, sucrose, lactose, fructose, maltose, molasses, optionally starch, optionally cellulose or from glycerol and ethanol. They are representatives of the Enterobacteriaceae family chosen from the genera Escherichia, Erwinia, Providencia and Serratia. The genera Escherichia and Serratia are preferred. Of the genus Escherichia the species Escherichia coli and of the genus Serratia the species Serratia marcescens are to be mentioned in particular.

25 30 Suitable strains, which produce L-threonine in particular, of the genus Escherichia, in particular of the species Escherichia coli. are, for example

Escherichia coli TF427
Escherichia coli H4578
Escherichia coli KY10935
Escherichia coli VNIIgenetika MG442
Escherichia coli VNIIgenetika M1
Escherichia coli VNIIgenetika 472T23
Escherichia coli BKIIM B-3996
Escherichia coli kat 13
Escherichia coli KCCM-10132

10 Suitable L-threonine-producing strains of the genus
Serratia, in particular of the species *Serratia marcescens*,
are, for example

Serratia marcescens HNr21
Serratia marcescens TLr156
Serratia marcescens T2000

15 Serratia marcescens T2000

Strains from the Enterobacteriaceae family which produce L-threonine preferably have, inter alia, one or more genetic or phenotypic features chosen from the group consisting of: resistance to α -amino- β -hydroxyvaleric acid, resistance to thialysine, resistance to ethionine, resistance to α -methylserine, resistance to diaminosuccinic acid, resistance to α -aminobutyric acid, resistance to borrelidin, resistance to rifampicin, resistance to valine analogues, such as, for example, valine hydroxamate, resistance to purine analogues, such as, for example, 6-dimethylaminopurine, a need for L-methionine, optionally a partial and compensable need for L-isoleucine, a need for meso-diaminopimelic acid, auxotrophy in respect of threonine-containing dipeptides, resistance to L-threonine, resistance to L-homoserine, resistance to L-lysine, resistance to L-methionine, resistance to L-glutamic acid, resistance to L-aspartate, resistance to L-leucine, resistance to L-phenylalanine, resistance to L-serine, resistance to L-cysteine, resistance to L-valine,

dehydrogenase, optionally an ability for sucrose utilization, enhancement of the threonine operon, enhancement of homoserine dehydrogenase I-aspartate kinase I, preferably of the feed back resistant form, enhancement 5 of homoserine kinase, enhancement of threonine synthase, enhancement of aspartate kinase, optionally of the feed back resistant form, enhancement of aspartate semialdehyde dehydrogenase, enhancement of phosphoenol pyruvate carboxylase, optionally of the feed back resistant form, 10 enhancement of phosphoenol pyruvate synthase, enhancement of transhydrogenase, enhancement of the RhtB gene product, enhancement of the RhtC gene product, enhancement of the YfiK gene product, enhancement of a pyruvate carboxylase, and attenuation of acetic acid formation.

15 It has been found that microorganisms of the Enterobacteriaceae family produce L-amino acids, in particular L-threonine, in an improved manner after enhancement, in particular over-expression, of at least one or more of the genes of the cysteine biosynthesis pathway 20 chosen from the group consisting of cysG, cysB, cysZ, cysK, cysM, cysA, cysW, cysU, cysP, cysD, cysN, cysC, cysJ, cysI, cysH, cysE and sbp.

The nucleotide sequences of the genes of *Escherichia coli* belong to the prior art (See following text references) and 25 can also be found in the genome sequence of *Escherichia coli* published by Blattner et al. (Science 277: 1453 - 1462 (1997)). The genes and activities of the cysteine biosynthesis pathway (cysteine biosynthetic pathway) are also described in summary form in Kredich (In: Neidhardt 30 (ed), *Escherichia coli and Salmonella*, American Society for Microbiology, Washington, D.C., USA: 514-527 (1996)).

cysG gene:

Description: Uroporphyrinogen III C methyl-transferase; uroporphyrin-2 oxidase; ferrochelatase

Reference: Peakman et al.; European Journal of Biochemistry 191(2): 315-323 (1990) Macdonald and Cole; Molecular and General Genetics 200(2): 328-334 (1985) Warren et al.; Biochemical Journal 265(3):725-729 (1990) Spencer et al.; FEBS Letters 335(1): 57-60 (1993)

Accession No.: AE000412

cysB gene:

10 Description: Positive regulator of the cys regulon,
transcription activator

Reference: Ostrowski et al.; Journal of Biological Chemistry 262(13): 5999-6005 (1987)
Mascarenhas and Yudkin; Molecular and General Genetics 177(3): 535-539 (1980)
Lochowska et al.; Journal of Biological Chemistry 276(3): 2098-2107 (2001)

Accession No.: AE000225

cysZ gene:

20 Description: Sulfate transporter
Reference: Byrne et al.; Journal of Bacteriology 152: 2153-2157 (1982)

Accession No.: AE000329

cysK gene:

25 Description: Cysteine synthase A, O-acetylserine
(thiol)-lyase A

EC No.:

Reference: Byrne et al.; Journal of Bacteriology 170(7): 3150-3157 (1988) Boronat et al.; Journal of General Microbiology 130: 673-685 (1984) Levy and Danchin; Molecular Microbiology 2(6): 777-783 (1988)

accession No.: AE000329

cysM gene:

Description: Cysteine synthase B, O-acetylserine
(thiol)-lyase B

EC No.: 4.2.99.8

5 Reference: Sirko et al.; Journal of Bacteriology 172(6): 3351-3357 (1990) Sirko et al.;
Journal of General Microbiology 133: 2719-2725 (1987)

Accession No.: AE000329

10 cysA gene:

Description: ATP-binding protein of the sulfate
transport system

Reference: Sirko et al.; Journal of Bacteriology 172(6): 3351-3357 (1990) Sirko et al.;
15 Journal of General Microbiology 133: 2719-2725 (1987)

Accession No.: AE000329

cysW gene:

Description: Membrane-bound sulfate transport protein

20 Reference: Sirko et al.; Journal of Bacteriology 172(6): 3351-3357 (1990)

Accession No.: AE000329, AE000330

cysU gene:

Description: Permease protein of the sulfate transport
system

25 Reference: Sirko et al.; Journal of Bacteriology 172(6): 3351-3357 (1990) Hryniwicz et al.;
Journal of Bacteriology 172(6): 3358-3366 (1990)

30 Accession No.: AE000330

Alternative gene name: cysT

cysP gene:

Description: Periplasmic thiosulfate-binding protein

Reference: Hrynniewicz et al.; Journal of Bacteriology 172(6): 3358-3366 (1990) Sirko et al.; Journal of Bacteriology 177(14): 4134-4136 (1995)

5 Accession No.: AE000330

cysD gene:

Description: Sub-unit 2 of ATP sulfurylase (ATP:sulfate adenylyl-transferase)

EC No.: 2.7.7.4

10 Reference: Leyh et al.; Journal of Biological Chemistry 267(15): 10405-10410 (1992) Leyh et al.; Journal of Biological Chemistry 263(5): 2409-2416 (1988)

Accession No.: AE000358

15 cysN gene:

Description: Sub-unit 1 of ATP sulfurylase (ATP:sulfate adenylyl-transferase)

EC No.: 2.7.7.4

Reference: Leyh et al.; Journal of Biological Chemistry 267(15): 10405-10410 (1992) Leyh et al.; Journal of Biological Chemistry 263(5): 2409-2416 (1988) Leyh and Suo; Journal of Biological Chemistry 267(1): 542-545 (1992)

20 25 Accession No.: AE000358

cysC gene:

Description: Adenylyl sulfate kinase (APS kinase)

EC No.: 2.7.1.25

Reference: Leyh et al.; Journal of Biological Chemistry 267(15): 10405-10410 (1992) Leyh et al.; Journal of Biological Chemistry 263(5): 2409-2416 (1988)

30 Accession No.: AE000358

cysJ gene:

Description: Flavoprotein of NADPH sulfite reductase
EC No.: 1.8.1.2
Reference: Ostrowski et al.; Journal of Biological
5 Chemistry 264(27): 15796-15808 (1989) Li et
al.; Gene 53(2-3): 227-234 (1987) Gaudu and
Fontecave; European Journal of Biochemistry
226(2): 459-463 (1994) Eschenbrenner et
al.; Journal of Biological Chemistry
10 270(35): 20550-20555 (1995)
Accession No.: AE000360
Alternative gene name: cysP

cysI gene:

Description: Haemoprotein of NADPH sulfite reductase
15 EC No.: 1.8.1.2
Reference: Ostrowski et al.; Journal of Biological
Chemistry 264(26): 15726-15737 (1989) Li et
al.; Gene 53(2-3): 227-234 (1987) Gaudu and
Fontecave; European Journal of Biochemistry
20 226(2): 459-463 (1994)
Accession No.: AE000360
Alternative gene name: cysQ

cysH gene:

Description: Phosphoadenosine phosphosulfate reductase
25 (PAPS reductase)
EC No.: 1.8.99.4
Reference: Ostrowski et al.; Journal of Biological
Chemistry 264(26): 15726-15737 (1989) Krone
et al.; Molecular and General Genetics
30 225(2): 314-319 (1991) Li et al.; Gene
53(2-3): 227-234 (1987) Berendt et al.;
European Journal of Biochemistry 233(1):
347-356 (1995)
Accession No.: AE000360

cysE gene:

Description: Serine acetyl-transferase
EC No.: 2.3.1.30
Reference: Denk and Böck; Journal of General
5 Microbiology 133, 515-25 (1987)
Accession No.: AE000438

sbp gene:

Description: Periplasmic sulfate-binding protein
Reference: Hellings and Evans, European Journal of
10 Biochemistry 149(2): 363-373 (1985) Sirkko
et al.; Journal of Bacteriology 177(14):
4134-4136 (1995) Jacobson et al.; Journal
of Biological Chemistry 266(8): 5220-5225
(1991)

15 Accession No.: AE000466

The nucleic acid sequences can be found in the databanks of
the National Center for Biotechnology Information (NCBI) of
the National Library of Medicine (Bethesda, MD, USA), the
nucleotide sequence databank of the European Molecular
20 Biologies Laboratories (EMBL, Heidelberg, Germany or
Cambridge, UK) or the DNA databank of Japan (DDBJ, Mishima,
Japan).

The genes described in the text references mentioned can be
used according to the invention. Alleles of the genes which
25 result from the degeneracy of the genetic code or due to
"sense mutations" of neutral function can furthermore be
used.

To achieve an enhancement, for example, expression of the
genes or the catalytic properties of the proteins can be
30 increased. The two measures can optionally be combined.

To achieve an over-expression, the number of copies of the
corresponding genes can be increased, or the promoter and
regulation region or the ribosome binding site upstream of

the structural gene can be mutated. Expression cassettes which are incorporated upstream of the structural gene act in the same way. By inducible promoters, it is additionally possible to increase the expression in the 5 course of fermentative L-threonine production. The expression is likewise improved by measures to prolong the life of the m-RNA. Furthermore, the enzyme activity is also enhanced by preventing the degradation of the enzyme protein. The genes or gene constructs can either be 10 present in plasmids with a varying number of copies, or can be integrated and amplified in the chromosome. Alternatively, an over-expression of the genes in question can furthermore be achieved by changing the composition of the media and the culture procedure.

15 Instructions in this context can be found by the expert, inter alia, in Chang and Cohen (Journal of Bacteriology 134: 1141-1156 (1978)), in Hartley and Gregori (Gene 13: 347-353 (1981)), in Amann and Brosius (Gene 40: 183-190 (1985)), in de Broer et al. (Proceedings of the National 20 Academy of Sciences of the United States of America 80: 21-25 (1983)), in LaVallie et al. (BIO/TECHNOLOGY 11: 187-193 (1993)), in PCT/US97/13359, in Llosa et al. (Plasmid 26: 222-224 (1991)), in Quandt and Klipp (Gene 80: 161-169 (1989)), in Hamilton (Journal of Bacteriology 171: 4617-4622 (1989)), in Jensen and Hammer (Biotechnology and Bioengineering 58: 191-195 (1998)) and in known textbooks 25 of genetics and molecular biology.

Plasmid vectors which are capable of replication in Enterobacteriaceae, such as e.g. cloning vectors derived 30 from pACYC184 (Bartolomé et al.; Gene 102: 75-78 (1991)), pTrc99A (Amann et al.; (Gene 69: 301-315 (1988)) or pSC101 derivatives (Vocke and Bastia, Proceedings of the National Academy of Sciences USA 80 (21): 6557-6561 (1983)) can be used. A strain transformed with a plasmid vector, where the 35 plasmid vector carries at least one or more of the genes

chosen from the group consisting of cysG, cysB, cysZ, cysK, cysM, cysA, cysW, cysU, cysP, cysD, cysN, cysC, cysJ, cysI, cysH, cysE and sbp or nucleotide sequences which code for these, can be employed in a process according to the
5 invention.

It is also possible to transfer mutations which affect the expression of the particular gene into various strains by sequence exchange (Hamilton et al. (Journal of Bacteriology 171: 4617 - 4622 (1989)), conjugation or transduction.

10 It may furthermore be advantageous for the production of L-amino acids, in particular L-threonine, with strains of the Enterobacteriaceae family to enhance one or more enzymes of the known threonine biosynthesis pathway or enzymes of anaplerotic metabolism or enzymes for the production of
15 reduced nicotinamide adenine dinucleotide phosphate or enzymes of glycolysis or PTS enzymes, in addition to enhancement of one or more of the genes of the cysteine biosynthesis pathway chosen from the group consisting of cysG, cysB, cysZ, cysK, cysM, cysA, cysW, cysU, cysP, cysD,
20 cysN, cysC, cysJ, cysI, cysH, cysE and sbp.

Thus, for example, at the same time one or more of the genes chosen from the group consisting of

- the thrABC operon which codes for aspartate kinase, homoserine dehydrogenase, homoserine kinase and threonine synthase (US-A-4,278,765),
25
- the pyc gene of Corynebacterium glutamicum which codes for pyruvate carboxylase (WO 99/18228),
- the pps gene which codes for phosphoenol pyruvate synthase (Molecular and General Genetics 231: 332-336
30 (1992)),
- the ppc gene which codes for phosphoenol pyruvate carboxylase (Gene 31: 279-283 (1984)),

- the pntA and pntB genes which code for transhydrogenase (European Journal of Biochemistry 158: 647-653 (1986)),
- the rhtB gene which imparts homoserine resistance (EP-A-0 994 190),

5 • the mqo gene which codes for malate:quinone oxidoreductase (WO 02/06459),

- the rhtC gene which imparts threonine resistance (EP-A-1 013 765),
- the thrE gene of *Corynebacterium glutamicum* which codes 10 for threonine export (WO 01/92545),
- the gdhA gene which codes for glutamate dehydrogenase (Nucleic Acids Research 11: 5257-5266 (1983); Gene 23: 199-209 (1983))
- the hns gene which codes for the DNA-binding protein 15 HLP-II (Molecular and General Genetics 212: 199-202 (1988)),
- the pgm gene which codes for phosphoglucomutase (Journal of Bacteriology 176: 5847-5851 (1994)),
- the fba gene which codes for fructose biphosphate 20 aldolase (Biochemical Journal 257: 529-534 (1989)),
- the ptsH gene of the ptsHIcrr operon which codes for the phosphohistidine protein hexose phosphotransferase of the phosphotransferase system PTS (Journal of Biological Chemistry 262: 16241-16253 (1987)),
- 25 • the ptsI gene of the ptsHIcrr operon which codes for enzyme I of the phosphotransferase system PTS (Journal of Biological Chemistry 262: 16241-16253 (1987)),
- the crr gene of the ptsHIcrr operon which codes for the glucose-specific IIA component of the phosphotransferase

system PTS (Journal of Biological Chemistry 262: 16241-16253 (1987)),

- the ptsG gene which codes for the glucose-specific IIBC component (Journal of Biological Chemistry 261: 16398-16403 (1986)),
5
- the lrp gene which codes for the regulator of the leucine regulon (Journal of Biological Chemistry 266: 10768-10774 (1991)),
10
- the mopB gene which codes for 10 Kd chaperone (Journal of Biological Chemistry 261: 12414-12419 (1986)) and is also known by the name groES,
15
- the ahpC gene of the ahpCF operon which codes for the small sub-unit of alkyl hydroperoxide reductase (Proceedings of the National Academy of Sciences USA 92: 7617-7621 (1995))
15
- the ahpF gene of the ahpCF operon which codes for the large sub-unit of alkyl hydroperoxide reductase (Proceedings of the National Academy of Sciences USA 92: 7617-7621 (1995))
20

can be enhanced, in particular over-expressed.

It may furthermore be advantageous for the production of L-amino acids, in particular L-threonine, in addition to enhancement of one or more of the genes chosen from the group consisting of cysG, cysB, cysZ, cysK, cysM, cysA, cysW, cysU, cysP, cysD, cysN, cysC, cysJ, cysI, cysH, cySE and sbp, for one or more of the genes chosen from the group consisting of
25

- the tdh gene which codes for threonine dehydrogenase (Ravnikar and Somerville (Journal of Bacteriology 169: 4716-4721 (1987))),
30

- the *mdh* gene which codes for malate dehydrogenase (E.C. 1.1.1.37) (Vogel et al. (Archives in Microbiology 149: 36-42 (1987))),
- the gene product of the open reading frame (orf) *yjfA* (Accession Number AAC77180 of the National Center for Biotechnology Information (NCBI, Bethesda, MD, USA)),
- the gene product of the open reading frame (orf) *ytfP* (Accession Number AAC77179 of the National Center for Biotechnology Information (NCBI, Bethesda, MD, USA)),
- the *pckA* gene which codes for the enzyme phosphoenol pyruvate carboxykinase (Medina et al. (Journal of Bacteriology 172: 7151-7156 (1990))),
- the *poxB* gene which codes for pyruvate oxidase (Grabau and Cronan (Nucleic Acids Research 14 (13), 5449-5460 (1986))),
- the *aceA* gene which codes for the enzyme isocitrate lyase (Matsuoko and McFadden (Journal of Bacteriology 170, 4528-4536 (1988))),
- the *dgsA* gene which codes for the DgsA regulator of the phosphotransferase system (Hosono et al. (Bioscience, Biotechnology and Biochemistry 59: 256-251 (1995)) and is also known under the name of the *mlc* gene,
- the *fruR* gene which codes for the fructose repressor (Jahreis et al. (Molecular and General Genetics 226: 332-336 (1991)) and is also known by the name of the *cra* gene, and
- the *rpoS* gene which codes for the sigma³⁸ factor (WO 01/05939) and is also known under the name of the *katF* gene,

30 to be attenuated, in particular eliminated or for the

The term "attenuation" in this connection describes the reduction or elimination of the intracellular activity or concentration of one or more enzymes or proteins in a microorganism which are coded by the corresponding DNA, for example by using a weak promoter or a gene or allele which codes for a corresponding enzyme or protein with a low activity or inactivates the corresponding enzyme or protein or gene, and optionally combining these measures.

By attenuation measures, the activity or concentration of the corresponding protein is in general reduced to 0 to 75%, 0 to 50%, 0 to 25%, 0 to 10% or 0 to 5% of the activity or concentration of the wild-type protein or of the activity or concentration of the protein in the starting microorganism.

15 It may furthermore be advantageous for the production of L-
amino acids, in particular L-threonine, in addition to
enhancement of one or more of the genes of the cysteine
biosynthesis pathway chosen from the group consisting of
cysG, cysB, cysZ, cysK, cysM, cysA, cysW, cysU, cysP, cysD,
20 cysN, cysC, cysJ, cysI, cysH, cysE and sbp, to eliminate
undesirable side reactions (Nakayama: "Breeding of Amino
Acid Producing Microorganisms", in: Overproduction of
Microbial Products, Krumphanzl, Sikyta, Vanek (eds.),
Academic Press, London, UK, 1982).

25 The microorganisms produced according to the invention can
be cultured in the batch process (batch culture), the fed
batch (feed process) or the repeated fed batch process
(repetitive feed process). A summary of known culture
methods is described in the textbook by Chmiel
30 (Bioprozesstechnik 1. Einführung in die
Bioverfahrenstechnik [Bioprocess Technology 1. Introduction
to Bioprocess Technology (Gustav Fischer Verlag, Stuttgart,
1991)) or in the textbook by Storhas (Bioreaktoren und
periphere Einrichtungen [Bioreactors and Peripheral

The culture medium to be used must meet the requirements of the particular strains in a suitable manner. Descriptions of culture media for various microorganisms are contained in the handbook "Manual of Methods for General 5 Bacteriology" of the American Society for Bacteriology (Washington D.C., USA, 1981).

Sugars and carbohydrates, such as e.g. glucose, sucrose, lactose, fructose, maltose, molasses, starch and optionally cellulose, oils and fats, such as e.g. soya oil, sunflower 10 oil, groundnut oil and coconut fat, fatty acids, such as e.g. palmitic acid, stearic acid and linoleic acid, alcohols, such as e.g. glycerol and ethanol, and organic acids, such as e.g. acetic acid, can be used as the source of carbon. These substances can be used individually or as 15 a mixture.

Organic nitrogen-containing compounds, such as peptones, yeast extract, meat extract, malt extract, corn steep liquor, soya bean flour and urea, or inorganic compounds, such as ammonium sulfate, ammonium chloride, ammonium 20 phosphate, ammonium carbonate and ammonium nitrate, can be used as the source of nitrogen. The sources of nitrogen can be used individually or as a mixture.

Phosphoric acid, potassium dihydrogen phosphate or dipotassium hydrogen phosphate or the corresponding sodium-25 containing salts can be used as the source of phosphorus. The culture medium must furthermore comprise salts of metals, such as e.g. magnesium sulfate or iron sulfate, which are necessary for growth. Finally, essential growth substances, such as amino acids and vitamins, can be 30 employed in addition to the above-mentioned substances. Suitable precursors can moreover be added to the culture medium. The starting substances mentioned can be added to the culture in the form of a single batch, or can be fed in during the culture in a suitable manner.

Basic compounds, such as sodium hydroxide, potassium hydroxide, ammonia or aqueous ammonia, or acid compounds, such as phosphoric acid or sulfuric acid, can be employed in a suitable manner to control the pH of the culture.

5 Antifoams, such as e.g. fatty acid polyglycol esters, can be employed to control the development of foam. Suitable substances having a selective action, e.g. antibiotics, can be added to the medium to maintain the stability of plasmids. To maintain aerobic conditions, oxygen or
10 oxygen-containing gas mixtures, such as e.g. air, are introduced into the culture. The temperature of the culture is usually 25°C to 45°C, and preferably 30°C to 40°C. Culturing is continued until a maximum of L-amino acids or L-threonine has formed. This target is usually
15 reached within 10 hours to 160 hours.

The analysis of L-amino acids can be carried out by anion exchange chromatography with subsequent ninhydrin derivation, as described by Spackman et al. (Analytical Chemistry 30: 1190-1206 (1958)), or it can take place by
20 reversed phase HPLC as described by Lindroth et al. (Analytical Chemistry 51: 1167-1174 (1979)).

The process according to the invention is used for the fermentative preparation of L-amino acids, such as, for example, L-threonine, L-isoleucine, L-valine, L-methionine,
25 L-homoserine and L-lysine, in particular L-threonine.

The minimal (M9) and complete media (LB) for Escherichia coli used are described by J.H. Miller (A short course in bacterial genetics (1992), Cold Spring Harbor Laboratory Press). The isolation of plasmid DNA from Escherichia coli
30 and all techniques of restriction, ligation, Klenow and alkaline phosphatase treatment are carried out by the method of Sambrook et al. (Molecular cloning - A Laboratory Manual (1989) Cold Spring Harbor Laboratory Press). Unless described otherwise, the transformation of Escherichia coli
35 is carried out by the method of Miller et al. (Principles

of the National Academy of Sciences of the United States of America (1989) 86: 2172-2175).

The incubation temperature for the preparation of strains and transformants is 37°C.

5 Example 1

Preparation of L-threonine using the cysB gene

1a) Construction of the expression plasmid pTrc99AcysB

The cysB gene from E. coli K12 is amplified using the polymerase chain reaction (PCR) and synthetic oligonucleotides. Starting from the nucleotide sequence of the cysB gene in E. coli K12 MG1655 (Accession Number AE000225, Blattner et al. (Science 277: 1453-1462 (1997))), PCR primers are synthesized (MWG Biotech, Ebersberg, Germany):

15 cysB1: 5' - GCGTCTAAGTGGATGGTTAAC - 3' (SEQ ID No. 1)

cysB2: 5' - GGTGCCGAAAATAACGCAAG - 3' (SEQ ID No. 2)

The chromosomal E. coli K12 MG1655 DNA employed for the PCR is isolated according to the manufacturer's instructions with "Qiagen Genomic-tips 100/G" (QIAGEN, Hilden, Germany). A DNA fragment approx. 1000 bp in size can be amplified with the specific primers under standard PCR conditions (Innis et al. (1990) PCR Protocols: A Guide to Methods and Applications, Academic Press) with Pfu-DNA polymerase (Promega Corporation, Madison, USA). The PCR product is ligated according to the manufacturer's instructions with the vector pCR-Blunt II-TOPO (Zero Blunt TOPO PCR Cloning Kit, Invitrogen, Groningen, The Netherlands) and transformed into the E. coli strain TOP10.

Selection of plasmid-carrying cells takes place on LB agar, to which 50 µg/ml kanamycin are added. After isolation of

cleaved with the restriction enzymes HindIII and XbaI and, after separation in 0.8% agarose gel, the cysB fragment is isolated with the aid of the QIAquick Gel Extraction Kit (QIAGEN, Hilden, Germany). The vector pTrc99A (Pharmacia Biotech, Uppsala, Sweden) is cleaved with the enzymes HindIII and XbaI and ligation is carried out with the cysB fragment isolated. The E. coli strain XL1-Blue MRF' (Stratagene, La Jolla, USA) is transformed with the ligation batch and plasmid-carrying cells are selected on LB agar, to which 50 µg/ml ampicillin are added. Successful cloning can be demonstrated after plasmid DNA isolation by control cleavage with the enzymes ScaI and SmaI. The plasmid is called pTrc99AcysB (Figure 1).

1b) Preparation of L-threonine with the strain
15 MG442/pTrc99AcysB

The L-threonine-producing E. coli strain MG442 is described in the patent specification US-A- 4,278,765 and deposited as CMIM B-1628 at the Russian National Collection for Industrial Microorganisms (VKPM, Moscow, Russia).

20 The strain MG442 is transformed with the expression plasmid pTrc99AcysB described in example Ia and with the vector pTrc99A and plasmid-carrying cells are selected on LB agar with 50 µg/ml ampicillin. The strains MG442/pTrc99AcysB and MG442/pTrc99A are formed in this manner. Selected
25 individual colonies are then multiplied further on minimal medium with the following composition: 3.5 g/l Na₂HPO₄*2H₂O, 1.5 g/l KH₂PO₄, 1 g/l NH₄Cl, 0.1 g/l MgSO₄*7H₂O, 2 g/l glucose, 20 g/l agar, 50 mg/l ampicillin. The formation of L-threonine is checked in batch cultures of 10 ml contained
30 in 100 ml conical flasks. For this, 10 ml of preculture medium of the following composition: 2 g/l yeast extract, 10 g/l (NH₄)₂SO₄, 1 g/l KH₂PO₄, 0.5 g/l MgSO₄*7H₂O, 15 g/l CaCO₃, 20 g/l glucose, 50 mg/l ampicillin are inoculated and the batch is incubated for 16 hours at 37°C and 180 rpm

on an ESR incubator from Kühner AG (Birsfelden, Switzerland).

250 µl portions of this preculture are transinoculated into 10 ml of production medium (25 g/l $(\text{NH}_4)_2\text{SO}_4$, 2 g/l KH_2PO_4 , 5 1 g/l $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, 0.03 g/l $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$, 0.018 g/l $\text{MnSO}_4 \cdot 1\text{H}_2\text{O}$, 30 g/l CaCO_3 , 20 g/l glucose, 50 mg/l ampicillin) and the batch is incubated for 48 hours at 37°C. For complete induction of the expression of the cysB gene, 100 mg/l isopropyl β-D-thiogalactopyranoside (IPTG) are added in 10 parallel batches. The formation of L-threonine by the starting strain MG442 is investigated in the same manner, but no addition of ampicillin to the medium takes place. After the incubation the optical density (OD) of the culture suspension is determined with an LP2W photometer 15 from Dr. Lange (Düsseldorf, Germany) at a measurement wavelength of 660 nm.

The concentration of L-threonine formed is then determined in the sterile-filtered culture supernatant with an amino acid analyzer from Eppendorf-BioTronik (Hamburg, Germany) 20 by ion exchange chromatography and post-column reaction with ninhydrin detection.

The result of the experiment is shown in Table 1.

Table 1

Strain	Additives	OD (660 nm)	L-Threonine g/l
MG442	-	5.6	1.4
MG442/pTrc99A	-	3.8	1.3
MG442/pTrc99AcysB	-	4.4	1.7
MG442/pTrc99AcysB	IPTG	5.4	2.0

Example 2

Preparation of L-threonine using the cysK gene

2a) Construction of the expression plasmid pTrc99AcysK

The cysK gene from E. coli K12 is amplified using the
5 polymerase chain reaction (PCR) and synthetic
oligonucleotides. Starting from the nucleotide sequence of
the cysK gene in E. coli K12 MG1655 (Accession Number
AE000329, Blattner et al. (Science 277: 1453-1462 (1997))),
PCR primers are synthesized (MWG Biotech, Ebersberg,

10 Germany):

cysK1: 5' - ·CAGTTAAGGACAGGCCATGAG - 3' (SEQ ID No. 3)

cysK2: 5' - GCTGGCATTACTGTTGCAATTC - 3' (SEQ ID No. 4)

The chromosomal E. coli K12 MG1655 DNA employed for the PCR
is isolated according to the manufacturer's instructions
15 with "Qiagen Genomic-tips 100/G" (QIAGEN, Hilden, Germany).
A DNA fragment approx. 1000 bp in size can be amplified
with the specific primers under standard PCR conditions
(Innis et al. (1990) PCR Protocols. A Guide to Methods and
Applications, Academic Press) with Pfu-DNA polymerase
20 (Promega Corporation, Madison, USA). The PCR product is
ligated according to the manufacturer's instructions with
the vector pCR-Blunt II-TOPO (Zero Blunt TOPO PCR Cloning
Kit, Invitrogen, Groningen, The Netherlands) and
transformed into the E. coli strain TOP10.

25 Selection of plasmid-carrying cells takes place on LB agar,
to which 50 µg/ml kanamycin are added. After isolation of
the plasmid DNA, the vector pCR-Blunt II-TOPO-cysK is
cleaved with the restriction enzymes SpeI and XbaI and,
after separation in 0.8% agarose gel, the cysK fragment is
30 isolated with the aid of the QIAquick Gel Extraction Kit
(QIAGEN, Hilden, Germany). The vector pTrc99A (Pharmacia
Biotech, Uppsala, Sweden) is cleaved with the enzyme XbaI

and ligation is carried out with the cysK fragment isolated. The E. coli strain XL1-Blue MRF' (Stratagene, La Jolla, USA) is transformed with the ligation batch and plasmid-carrying cells are selected on LB agar, to which 5 50 µg/ml ampicillin are added. Successful cloning can be demonstrated after plasmid DNA isolation by control cleavage with the enzymes HindIII and PvuII. The plasmid is called pTrc99AcysK (Figure 2).

2b) Preparation of L-threonine with the strain
10 MG442/pTrc99AcysK

The L-threonine-producing E. coli strain MG442 is described in the patent specification US-A- 4,278,765 and deposited as CMIM B-1628 at the Russian National Collection for Industrial Microorganisms (VKPM, Moscow, Russia).

15 The strain MG442 is transformed with the expression plasmid pTrc99AcysK described in example 2a and with the vector pTrc99A and plasmid-carrying cells are selected on LB agar with 50 µg/ml ampicillin. The strains MG442/pTrc99AcysK and MG442/pTrc99A are formed in this manner. Selected
20 individual colonies are then multiplied further on minimal medium with the following composition: 3.5 g/l Na₂HPO₄*2H₂O, 1.5 g/l KH₂PO₄, 1 g/l NH₄Cl, 0.1 g/l MgSO₄*7H₂O, 2 g/l glucose, 20 g/l agar, 50 mg/l ampicillin. The formation of L-threonine is checked in batch cultures of 10 ml contained
25 in 100 ml conical flasks. For this, 10 ml of preculture medium of the following composition: 2 g/l yeast extract, 10 g/l (NH₄)₂SO₄, 1 g/l KH₂PO₄, 0.5 g/l MgSO₄*7H₂O, 15 g/l CaCO₃, 20 g/l glucose, 50 mg/l ampicillin are inoculated and the batch is incubated for 16 hours at 37°C and 180 rpm
30 on an ESR incubator from Kühner AG (Birsfelden, Switzerland).

250 µl portions of this preculture are transinoculated into 10 ml of production medium (25 g/l (NH₄)₂SO₄, 2 g/l KH₂PO₄, 1 g/l yeast extract, 0.5 g/l MgSO₄*7H₂O, 15 g/l CaCO₃, 20 g/l glucose, 50 mg/l ampicillin, 0.005 g/l MnSO₄*1H₂O,

30 g/l CaCO₃, 20 g/l glucose, 50 mg/l ampicillin) and the batch is incubated for 48 hours at 37°C. The formation of L-threonine by the starting strain MG442 is investigated in the same manner, but no addition of ampicillin to the 5 medium takes place. After the incubation the optical density (OD) of the culture suspension is determined with an LP2W photometer from Dr. Lange (Düsseldorf, Germany) at a measurement wavelength of 660 nm.

The concentration of L-threonine formed is then determined 10 in the sterile-filtered culture supernatant with an amino acid analyzer from Eppendorf-BioTronik (Hamburg, Germany) by ion exchange chromatography and post-column reaction with ninhydrin detection.

The result of the experiment is shown in Table 2.

15

Table 2

Strain	OD (660 nm)	L-Threonine g/l
MG442	5.6	1.4
MG442/pTrc99A	3.8	1.3
MG442/pTrc99AcysK	5.6	2.1

Example 3

Preparation of L-threonine using the cysM gene

3a) Construction of the expression plasmid pTrc99AcysM

20 The cysM gene from *E. coli* K12 is amplified using the polymerase chain reaction (PCR) and synthetic oligonucleotides. Starting from the nucleotide sequence of the cysM gene in *E. coli* K12 NCBI accession number

AE000329, Blattner et al. (Science 277: 1453-1462 (1997)), PCR primers are synthesized (MWG Biotech, Ebersberg, Germany). The sequences of the primers are modified such that recognition sites for restriction enzymes are formed.

5 The recognition sequence for XbaI is chosen for the cysM1 primer and the recognition sequence for HindIII for the cysM2 primer, which are marked by underlining in the nucleotide sequence shown below:

cysM1: 5' - CGCATCAGTCTAGACCACGTTAGGATAG - 3'
10 (SEQ ID No. 5)

cysM2: 5' - CATCAGTCTCCGAAGCTTTAATCC - 3'
(SEQ ID No. 6)

The chromosomal E. coli K12 MG1655 DNA employed for the PCR is isolated according to the manufacturer's instructions
15 with "Qiagen Genomic-tips 100/G" (QIAGEN, Hilden, Germany). A DNA fragment approx. 950 bp in size can be amplified with the specific primers under standard PCR conditions (Innis et al. (1990) PCR Protocols. A Guide to Methods and Applications, Academic Press) with Pfu-DNA polymerase
20 (Promega Corporation, Madison, USA). The PCR product is ligated according to the manufacturer's instructions with the vector pCR-Blunt II-TOPO (Zero Blunt TOPO PCR Cloning Kit, Invitrogen, Groningen, The Netherlands) and transformed into the E. coli strain TOP10.
25 Selection of plasmid-carrying cells takes place on LB agar, to which 50 µg/ml kanamycin are added. After isolation of the plasmid DNA, the vector pCR-Blunt II-TOPO-cysM is cleaved with the restriction enzymes HindIII and XbaI and, after separation in 0.8% agarose gel, the cysM fragment is
30 isolated with the aid of the QIAquick Gel Extraction Kit (QIAGEN, Hilden, Germany). The vector pTrc99A (Pharmacia Biotech, Uppsala, Sweden) is cleaved with the enzymes HindIII and XbaI and ligation is carried out with the cysM

(Stratagene, La Jolla, USA) is transformed with the ligation batch and plasmid-carrying cells are selected on LB agar, to which 50 µg/ml ampicillin are added. Successful cloning can be demonstrated after plasmid DNA isolation by 5 control cleavage with the enzymes EcoRV, Eco91I, PauI and SspI. The plasmid is called pTrc99AcysM (Figure 3).

3b) Preparation of L-threonine with the strain
MG442/pTrc99AcysM

The L-threonine-producing E. coli strain MG442 is described 10 in the patent specification US-A- 4,278,765 and deposited as CMIM B-1628 at the Russian National Collection for Industrial Microorganisms (VKPM, Moscow, Russia).

The strain MG442 is transformed with the expression plasmid pTrc99AcysM described in example 3a and with the vector 15 pTrc99A and plasmid-carrying cells are selected on LB agar with 50 µg/ml ampicillin. The strains MG442/pTrc99AcysM and MG442/pTrc99A are formed in this manner. Selected individual colonies are then multiplied further on minimal medium with the following composition: 3.5 g/l Na₂HPO₄*2H₂O, 20 1.5 g/l KH₂PO₄, 1 g/l NH₄Cl, 0.1 g/l MgSO₄*7H₂O, 2 g/l glucose, 20 g/l agar, 50 mg/l ampicillin. The formation of L-threonine is checked in batch cultures of 10 ml contained in 100 ml conical flasks. For this, 10 ml of preculture medium of the following composition: 2 g/l yeast extract, 25 10 g/l (NH₄)₂SO₄, 1 g/l KH₂PO₄, 0.5 g/l MgSO₄*7H₂O, 15 g/l CaCO₃, 20 g/l glucose, 50 mg/l ampicillin are inoculated and the batch is incubated for 16 hours at 37°C and 180 rpm on an ESR incubator from Kühner AG (Birsfelden, Switzerland).
30 250 µl portions of this preculture are transinoculated into 10 ml of production medium (25 g/l (NH₄)₂SO₄, 2 g/l KH₂PO₄, 1 g/l MgSO₄*7H₂O, 0.03 g/l FeSO₄*7H₂O, 0.018 g/l MnSO₄*1H₂O, 30 g/l CaCO₃, 20 g/l glucose, 50 mg/l ampicillin) and the batch is incubated for 48 hours at 37°C. The formation of

L-threonine by the starting strain MG442 is investigated in the same manner, but no addition of ampicillin to the medium takes place. After the incubation the optical density (OD) of the culture suspension is determined with 5 an LP2W photometer from Dr. Lange (Düsseldorf, Germany) at a measurement wavelength of 660 nm.

The concentration of L-threonine formed is then determined in the sterile-filtered culture supernatant with an amino acid analyzer from Eppendorf-BioTronik (Hamburg, Germany) 10 by ion exchange chromatography and post-column reaction with ninhydrin detection.

The result of the experiment is shown in Table 3.

Table 3

Strain	OD (660 nm)	L-Threonine g/l
MG442	5.6	1.4
MG442/pTrc99A	3.8	1.3
MG442/pTrc99AcysM	1.6	2.0

15 Example 4

Preparation of L-threonine using the cysP, cysU, cysW and cysA genes

4a) Construction of the expression plasmid pTrc99AcysPUWA

The cysP, cysU, cysW and cysA genes from *E. coli* K12 are 20 amplified using the polymerase chain reaction (PCR) and synthetic oligonucleotides. Starting from the nucleotide sequence of the cysP, cysU, cysW and cysA genes in *E. coli* K12 MG1655 (Accession Number AE000329 and AE000330,

Blattner et al. (Science 277: 1453-1462 (1997)), PCR primers are synthesized (MWG Biotech, Ebersberg, Germany). The sequences of the primers are modified such that recognition sites for restriction enzymes are formed. The 5 recognition sequence for XbaI is chosen for the cysPUWA1 primer and the recognition sequence for HindIII for the cysPUWA2 primer, which are marked by underlining in the nucleotide sequence shown below:

cysPUWA1: 5' - GTCTCTAGATAAAATAAGGGTGCGCAATGGC - 3'
10 (SEQ ID No. 7)

cysPUWA2: 5' - CCGGGCGTTTAAGCTTCACTCAACC - 3'
(SEQ ID No. 8)

The chromosomal E. coli K12 MG1655 DNA employed for the PCR is isolated according to the manufacturer's instructions 15 with "Qiagen Genomic-tips 100/G" (QIAGEN, Hilden, Germany). A DNA fragment approx. 3900 bp in size can be amplified with the specific primers under standard PCR conditions (Innis et al. (1990) PCR Protocols. A Guide to Methods and Applications, Academic Press) with Pfu-DNA polymerase 20 (Promega Corporation, Madison, USA). The PCR product is cleaved with the restriction enzymes XbaI and HindIII and ligated with the vector pTrc99A (Pharmacia Biotech, Uppsala, Sweden), which has been digested with the enzymes XbaI and HindIII. The E. coli strain XL1-Blue MRF' 25 (Stratagene, La Jolla, USA) is transformed with the ligation batch and plasmid-carrying cells are selected on LB agar, to which 50 µg/ml ampicillin are added. Successful cloning can be demonstrated after plasmid DNA isolation by control cleavage with the enzymes BamHI, EcoRV, MluI, NdeI 30 and SspI. The plasmid is called pTrc99AcysPUWA (Figure 4).

4b) Preparation of L-threonine with the strain
MG442/pTrc99AcysPUWA

The L-threonine-producing *E. coli* strain MG442 is described in the patent specification US-A- 4,278,765 and deposited 5 as CMIM B-1628 at the Russian National Collection for Industrial Microorganisms (VKPM, Moscow, Russia).

The strain MG442 is transformed with the expression plasmid pTrc99AcysPUWA described in example 4a and with the vector pTrc99A and plasmid-carrying cells are selected on LB agar 10 with 50 µg/ml ampicillin. The strains MG442/pTrc99AcysPUWA and MG442/pTrc99A are formed in this manner. Selected individual colonies are then multiplied further on minimal medium with the following composition: 3.5 g/l Na₂HPO₄*2H₂O, 1.5 g/l KH₂PO₄, 1 g/l NH₄Cl, 0.1 g/l MgSO₄*7H₂O, 2 g/l 15 glucose, 20 g/l agar, 50 mg/l ampicillin. The formation of L-threonine is checked in batch cultures of 10 ml contained in 100 ml conical flasks. For this, 10 ml of preculture medium of the following composition: 2 g/l yeast extract, 10 g/l (NH₄)₂SO₄, 1 g/l KH₂PO₄, 0.5 g/l MgSO₄*7H₂O, 15 g/l 20 CaCO₃, 20 g/l glucose, 50 mg/l ampicillin are inoculated and the batch is incubated for 16 hours at 37°C and 180 rpm on an ESR incubator from Kühner AG (Birsfelden, Switzerland).

250 µl portions of this preculture are transinoculated into 25 10 ml of production medium (25 g/l (NH₄)₂SO₄, 2 g/l KH₂PO₄, 1 g/l MgSO₄*7H₂O, 0.03 g/l FeSO₄*7H₂O, 0.018 g/l MnSO₄*1H₂O, 30 g/l CaCO₃, 20 g/l glucose, 50 mg/l ampicillin) and the batch is incubated for 48 hours at 37°C. For complete induction of the expression of the cysPUWA genes, 100 mg/l 30 isopropyl β-D-thiogalactopyranoside (IPTG) are added in parallel batches. The formation of L-threonine by the starting strain MG442 is investigated in the same manner, but no addition of ampicillin to the medium takes place. After the incubation the optical density (OD) of the 35 culture is measured in Miller and Miller units.

from Dr. Lange (Düsseldorf, Germany) at a measurement wavelength of 660 nm.

The concentration of L-threonine formed is then determined in the sterile-filtered culture supernatant with an amino acid analyzer from Eppendorf-BioTronik (Hamburg, Germany) by ion exchange chromatography and post-column reaction with ninhydrin detection.

The result of the experiment is shown in Table 4.

Table 4

Strain	Additives	OD (660 nm)	L-Threonine g/l
MG442	-	5.6	1.4
MG442/pTrc99A	-	3.8	1.3
MG442/pTrc99AcysPUWA	-	5.5	1.7
MG442/pTrc99AcysPUWA	IPTG	6.5	2.1

10

Example 5

Preparation of L-threonine using the cysD, cysN and cysC genes

5a) Construction of the expression plasmid pTrc99AcysDNC

15 The cysD, cysN and cysC genes from *E. coli* K12 are amplified using the polymerase chain reaction (PCR) and synthetic oligonucleotides. Starting from the nucleotide sequence of the cysD, cysN and cysC genes in *E. coli* K12 MG1655 (Accession Number AE000358, Blattner et al. (Science 20 277: 1453-1462 (1997))), PCR primers are synthesized (MWG Biotech, Ebersberg, Germany). The sequences of the primers

are modified such that recognition sites for restriction enzymes are formed. The recognition sequence for XbaI is chosen for the cysDNC1 primer and the recognition sequence for HindIII for the cysDNC2 primer, which are marked by underlining in the nucleotide sequence shown below:

5 cysDNC1: 5' - GCAAGAAAATAGCGGTCTAGATAAGGAACG - 3'

(SEQ ID No. 9)

cysDNC2: 5' - CATGGAAAGCTTGTGGTGTCTCAGG - 3'

(SEQ ID No. 10)

10 The chromosomal E. coli K12 MG1655 DNA employed for the PCR is isolated according to the manufacturer's instructions with "Qiagen Genomic-tips 100/G" (QIAGEN, Hilden, Germany). A DNA fragment approx. 3000 bp in size can be amplified with the specific primers under standard PCR conditions
15 (Innis et al. (1990) PCR Protocols. A Guide to Methods and Applications, Academic Press) with Pfu-DNA polymerase (Promega Corporation, Madison, USA). The PCR product is cleaved with the restriction enzymes XbaI and HindIII and ligated with the vector pTrc99A (Pharmacia Biotech, Uppsala, Sweden), which has been digested with the enzymes XbaI and HindIII. The E. coli strain XL1-Blue MRF'
20 (Stratagene, La Jolla, USA) is transformed with the ligation batch and plasmid-carrying cells are selected on LB agar, to which 50 µg/ml ampicillin are added. Successful cloning can be demonstrated after plasmid DNA isolation by
25 control cleavage with the enzymes EcoRV, HincII, NruI, PvuI and ScaI. The plasmid is called pTrc99AcysDNC (Figure 5).

5b) Preparation of L-threonine with the strain
MG442/pTrc99AcysDNC

30 The L-threonine-producing E. coli strain MG442 is described in the patent specification US-A- 4,278,765 and deposited as CMIM B-1628 at the Russian National Collection for Industrial Microorganisms (VKPM, Moscow, Russia).

The strain MG442 is transformed with the expression plasmid pTrc99AcysDNC described in example 5a and with the vector pTrc99A and plasmid-carrying cells are selected on LB agar with 50 µg/ml ampicillin. The strains MG442/pTrc99AcysDNC 5 and MG442/pTrc99A are formed in this manner. Selected individual colonies are then multiplied further on minimal medium with the following composition: 3.5 g/l Na₂HPO₄*2H₂O, 1.5 g/l KH₂PO₄, 1 g/l NH₄Cl, 0.1 g/l MgSO₄*7H₂O, 2 g/l glucose, 20 g/l agar, 50 mg/l ampicillin. The formation of 10 L-threonine is checked in batch cultures of 10 ml contained in 100 ml conical flasks. For this, 10 ml of preculture medium of the following composition: 2 g/l yeast extract, 10 g/l (NH₄)₂SO₄, 1 g/l KH₂PO₄, 0.5 g/l MgSO₄*7H₂O, 15 g/l CaCO₃, 20 g/l glucose, 50 mg/l ampicillin are inoculated 15 and the batch is incubated for 16 hours at 37°C and 180 rpm on an ESR incubator from Kühner AG (Birsfelden, Switzerland).

250 µl portions of this preculture are transinoculated into 10 ml of production medium (25 g/l (NH₄)₂SO₄, 2 g/l KH₂PO₄, 20 1 g/l MgSO₄*7H₂O, 0.03 g/l FeSO₄*7H₂O, 0.018 g/l MnSO₄*1H₂O, 30 30 g/l CaCO₃, 20 g/l glucose, 50 mg/l ampicillin) and the batch is incubated for 48 hours at 37°C. The formation of L-threonine by the starting strain MG442 is investigated in the same manner, but no addition of ampicillin to the 25 medium takes place. After the incubation the optical density (OD) of the culture suspension is determined with an LP2W photometer from Dr. Lange (Düsseldorf, Germany) at a measurement wavelength of 660 nm.

30 The concentration of L-threonine formed is then determined in the sterile-filtered culture supernatant with an amino acid analyzer from Eppendorf-BioTronik (Hamburg, Germany) by ion exchange chromatography and post-column reaction with ninhydrin detection.

The result of the experiment is shown in Table 5.

Table 5

Strain	OD (660 nm)	L-Threonine g/l
MG442	5.6	1.4
MG442/pTrc99A	3.8	1.3
MG442/pTrc99AcysDNC	5.1	2.5

Example 6

Preparation of L-threonine using the cysJ and cysI genes

5 6a) Construction of the expression plasmid pTrc99AcysJI

The cysJ and cysI genes from E. coli K12 are amplified using the polymerase chain reaction (PCR) and synthetic oligonucleotides. Starting from the nucleotide sequence of the cysJ and cysI genes in E. coli K12 MG1655 (Accession 10 Number AE000360, Blattner et al. (Science 277: 1453-1462 (1997)), PCR primers are synthesized (MWG Biotech, Ebersberg, Germany):

cysJI1: 5' - CTGGAACATAACCGACGCATGAC - 3' (SEQ ID No. 11)

cysJI2: 5' - GACCGGGCTGATGGTTAATCC - 3' (SEQ ID No. 12)

15 The chromosomal E. coli K12 MG1655 DNA employed for the PCR is isolated according to the manufacturer's instructions with "Qiagen Genomic-tips 100/G" (QIAGEN, Hilden, Germany). A DNA fragment approx. 3550 bp in size can be amplified with the specific primers under standard PCR conditions 20 (Innis et al. (1990) PCR Protocols. A Guide to Methods and Applications, Academic Press) with Pfu-DNA polymerase (Promega Corporation, Madison, USA). The PCR product is ligated according to the manufacturer's instructions with

the vector pCR-Blunt II-TOPO (Zero Blunt TOPO PCR Cloning Kit, Invitrogen, Groningen, The Netherlands) and transformed into the *E. coli* strain TOP10.

Selection of plasmid-carrying cells takes place on LB agar, 5 to which 50 µg/ml kanamycin are added. After isolation of the plasmid DNA, the vector pCR-Blunt II-TOPO-cysJI is cleaved with the restriction enzymes HindIII and XbaI and, after separation in 0.8% agarose gel, the cysJI fragment is isolated with the aid of the QIAquick Gel Extraction Kit 10 (QIAGEN, Hilden, Germany). The vector pTrc99A (Pharmacia Biotech, Uppsala, Sweden) is cleaved with the enzymes HindIII and XbaI and ligation is carried out with the cysJI fragment isolated. The *E. coli* strain XL1-Blue MRF' 15 (Stratagene, La Jolla, USA) is transformed with the ligation batch and plasmid-carrying cells are selected on LB agar, to which 50 µg/ml ampicillin are added. Successful cloning can be demonstrated after plasmid DNA isolation by control cleavage with the enzymes AccI, ClaI and SphI. The plasmid is called pTrc99AcysJI (Figure 6).

20 6b) Preparation of L-threonine with the strain
MG442/pTrc99AcysJI

The L-threonine-producing *E. coli* strain MG442 is described in the patent specification US-A- 4,278,765 and deposited as CMIM B-1628 at the Russian National Collection for 25 Industrial Microorganisms (VKPM, Moscow, Russia).

The strain MG442 is transformed with the expression plasmid pTrc99AcysJI described in example 6a and with the vector pTrc99A and plasmid-carrying cells are selected on LB agar with 50 µg/ml ampicillin. The strains MG442/pTrc99AcysJI 30 and MG442/pTrc99A are formed in this manner. Selected individual colonies are then multiplied further on minimal medium with the following composition: 3.5 g/l Na₂HPO₄*2H₂O, 1.5 g/l KH₂PO₄, 1 g/l NH₄Cl, 0.1 g/l MgSO₄*7H₂O, 2 g/l glucose, 20 g/l agar, 50 µg/l ampicillin. The formation of

L-threonine is checked in batch cultures of 10 ml contained in 100 ml conical flasks. For this, 10 ml of preculture medium of the following composition: 2 g/l yeast extract, 10 g/l $(\text{NH}_4)_2\text{SO}_4$, 1 g/l KH_2PO_4 , 0.5 g/l $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, 15 g/l 5 CaCO_3 , 20 g/l glucose, 50 mg/l ampicillin are inoculated and the batch is incubated for 16 hours at 37°C and 180 rpm on an ESR incubator from Kühner AG (Birsfelden, Switzerland).

250 μl portions of this preculture are transinoculated into
10 10 ml of production medium (25 g/l $(\text{NH}_4)_2\text{SO}_4$, 2 g/l KH_2PO_4 , 1 g/l $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, 0.03 g/l $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$, 0.018 g/l $\text{MnSO}_4 \cdot 1\text{H}_2\text{O}$, 30 g/l CaCO_3 , 20 g/l glucose, 50 mg/l ampicillin) and the batch is incubated for 48 hours at 37°C. The formation of L-threonine by the starting strain MG442 is investigated in
15 the same manner, but no addition of ampicillin to the medium takes place. After the incubation the optical density (OD) of the culture suspension is determined with an LP2W photometer from Dr. Lange (Düsseldorf, Germany) at a measurement wavelength of 660 nm.
20 The concentration of L-threonine formed is then determined in the sterile-filtered culture supernatant with an amino acid analyzer from Eppendorf-BioTronik (Hamburg, Germany) by ion exchange chromatography and post-column reaction with ninhydrin detection.
25 The result of the experiment is shown in Table 6.

Table 6

Strain	OD (660 nm)	L-Threonine g/l
MG442	5.6	1.4
MG442/pTrc99A	3.8	1.3
MG442/pTrc99AcysJI	6.3	2.5

Example 7

Preparation of L-threonine using the cysH gene

5 7a) Construction of the expression plasmid pTrc99AcysH

The cysH gene from E. coli K12 is amplified using the polymerase chain reaction (PCR) and synthetic oligonucleotides. Starting from the nucleotide sequence of the cysH gene in E. coli K12 MG1655 (Accession Number AE000360, Blattner et al. (Science 277: 1453-1462 (1997))), PCR primers are synthesized (MWG Biotech, Ebersberg, Germany):

cysH1: 5' - GGCAAACAGTGAGGAATCTATG - 3' (SEQ ID No. 13)

cysH2: 5' - GTCCGGCAATATTACCCCTTC - 3' (SEQ ID No. 14)

15 The chromosomal E. coli K12 MG1655 DNA employed for the PCR is isolated according to the manufacturer's instructions with "Qiagen Genomic-tips 100/G" (QIAGEN, Hilden, Germany). A DNA fragment approx. 800 bp in size can be amplified with the specific primers under standard PCR conditions (Innis et al. (1990) PCR Protocols. A Guide to Methods and Applications, Academic Press) with Pfu-DNA polymerase (Promega Corporation, Madison, USA). The PCR product is digested according to the manufacturer's instructions with

the vector pCR-Blunt II-TOPO (Zero Blunt TOPO PCR Cloning Kit, Invitrogen, Groningen, The Netherlands) and transformed into the E. coli strain TOP10.

Selection of plasmid-carrying cells takes place on LB agar, 5 to which 50 µg/ml kanamycin are added. After isolation of the plasmid DNA, the vector pCR-Blunt II-TOPO-cysH is cleaved with the restriction enzymes HindIII and XbaI and, after separation in 0.8% agarose gel, the cysH fragment is isolated with the aid of the QIAquick Gel Extraction Kit 10 (QIAGEN, Hilden, Germany). The vector pTrc99A (Pharmacia Biotech, Uppsala, Sweden) is cleaved with the enzymes HindIII and XbaI and ligation is carried out with the cysH fragment isolated. The E. coli strain XL1-Blue MRF' 15 (Stratagene, La Jolla, USA) is transformed with the ligation batch and plasmid-carrying cells are selected on LB agar, to which 50 µg/ml ampicillin are added. Successful cloning can be demonstrated after plasmid DNA isolation by control cleavage with the enzymes HincII and MluI. The plasmid is called pTrc99AcysH (Figure 7).

20 7b) Preparation of L-threonine with the strain
MG442/pTrc99AcysH

The L-threonine-producing E. coli strain MG442 is described in the patent specification US-A- 4,278,765 and deposited as CMIM B-1628 at the Russian National Collection for 25 Industrial Microorganisms (VKPM, Moscow, Russia).

The strain MG442 is transformed with the expression plasmid pTrc99AcysH described in example 7a and with the vector pTrc99A and plasmid-carrying cells are selected on LB agar with 50 µg/ml ampicillin. The strains MG442/pTrc99AcysH and 30 MG442/pTrc99A are formed in this manner. Selected individual colonies are then multiplied further on minimal medium with the following composition: 3.5 g/l Na₂HPO₄*2H₂O, 1.5 g/l KH₂PO₄, 1 g/l NH₄Cl, 0.1 g/l MgSO₄*7H₂O, 2 g/l yeast extract, 0.1 g/l glucose, 50 µg/ml ampicillin. The formation of

L-threonine is checked in batch cultures of 10 ml contained in 100 ml conical flasks. For this, 10 ml of preculture medium of the following composition: 2 g/l yeast extract, 10 g/l $(\text{NH}_4)_2\text{SO}_4$, 1 g/l KH_2PO_4 , 0.5 g/l $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, 15 g/l 5 CaCO_3 , 20 g/l glucose, 50 mg/l ampicillin are inoculated and the batch is incubated for 16 hours at 37°C and 180 rpm on an ESR incubator from Kühner AG (Birsfelden, Switzerland).

250 μl portions of this preculture are transinoculated into 10 10 ml of production medium (25 g/l $(\text{NH}_4)_2\text{SO}_4$, 2 g/l KH_2PO_4 , 1 g/l $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, 0.03 g/l $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$, 0.018 g/l $\text{MnSO}_4 \cdot 1\text{H}_2\text{O}$, 30 g/l CaCO_3 , 20 g/l glucose, 50 mg/l ampicillin) and the batch is incubated for 48 hours at 37°C. The formation of L-threonine by the starting strain MG442 is investigated in 15 the same manner, but no addition of ampicillin to the medium takes place. After the incubation the optical density (OD) of the culture suspension is determined with an LP2W photometer from Dr. Lange (Düsseldorf, Germany) at a measurement wavelength of 660 nm.

20 The concentration of L-threonine formed is then determined in the sterile-filtered culture supernatant with an amino acid analyzer from Eppendorf-BioTronik (Hamburg, Germany) by ion exchange chromatography and post-column reaction with ninhydrin detection.

25 The result of the experiment is shown in Table 7.

Table 7

Strain	OD (660 nm)	L-Threonine g/l
MG442	5.6	1.4
MG442/pTrc99A	3.8	1.3
MG442/pTrc99AcysH	4.1	2.7

Brief Description of the Figures:

Figure 1: Map of the plasmid pTrc99AcysB containing the
5 cysB gene.

Figure 2: Map of the plasmid pTrc99AcysK containing the
cysK gene.

Figure 3: Map of the plasmid pTrc99AcysM containing the
cysM gene.

10 Figure 4: Map of the plasmid pTrc99AcysPUWA containing
the cysP, cysU, cysW and cysA genes.

Figure 5: Map of the plasmid pTrc99AcysDNC containing the
cysD, cysN and cysC genes.

15 Figure 6: Map of the plasmid pTrc99AcysJI containing the
cysJ and cysI genes.

Figure 7: Map of the plasmid pTrc99AcysH containing the
cysH gene.

The length data are to be understood as approx. data. The
abbreviations and designations used have the following
20 meaning:

: --- : Repetitive nucleotide sequence

- lacI: Gene for the repressor protein of the trc promoter
- Ptrc: trc promoter region, IPTG-inducible
- cysB: Coding region of the cysB gene
- 5 • cysK: Coding region of the cysK gene
- cysM: Coding region of the cysM gene
- cysP: Coding region of the cysP gene
- cysU: Coding region of the cysU gene
- cysW: Coding region of the cysW gene
- 10 • cysA: Coding region of the cysA gene
- cysD: Coding region of the cysD gene
- cysN: Coding region of the cysN gene
- cysC: Coding region of the cysC gene
- cysJ: Coding region of the cysJ gene
- 15 • cysI: Coding region of the cysI gene
- cysH: Coding region of the cysH gene
- 5S: 5S rRNA region
- rrnBT: rRNA terminator region

The abbreviations for the restriction enzymes have the
20 following meaning .

- AccI: Restriction endonuclease from *Acinetobacter calcoaceticus*
- BamHI: Restriction endonuclease from *Bacillus*

- *Bst*EII: Restriction endonuclease from *Bacillus stearothermophilus* ATCC 12980
- *Cla*I: Restriction endonuclease from *Caryophannion latum*
- *Eco*RI: Restriction endonuclease from *Escherichia coli*

5 RY13

- *Eco*RV: Restriction endonuclease from *Escherichia coli* B946
- *Hinc*II: Restriction endonuclease from *Haemophilus influenzae* R_c

10 • *Hind*III: Restriction endonuclease from *Haemophilus influenzae*

- *Mlu*I: Restriction endonuclease from *Micrococcus luteus* IFO 12992
- *Nde*I: Restriction endonuclease from *Neisseria dentrificans*

15 • *Nru*I: Restriction endonuclease from *Norcadia ruba* (ATCC 15906)

- *Pau*I: Restriction endonuclease from *Paracoccus alcaliphilus*

20 • *Pvu*I: Restriction endonuclease from *Proteus vulgaris* (ATCC 13315)

- *Pvu*II: Restriction endonuclease from *Proteus vulgaris* (ATCC 13315)
- *Sca*I: Restriction endonuclease from *Streptomyces caespitosus*

25 • *Sma*I: Restriction endonuclease from *Serratia marcescens*

- SpeI: Restriction endonuclease from *Sphaerotilus* species ATCC 13923
- SphI: Restriction endonuclease from *Streptomyces phaeochromogenes*
- 5 • SspI: Restriction endonuclease from *Sphaerotilus* species ATCC 13925
- XbaI: Restriction endonuclease from *Xanthomonas campestris*

What is claimed is:

1. A process for the preparation of L-amino acids, in particular L-threonine, which comprises carrying out the following steps:
 - 5 a) fermentation of microorganisms of the Enterobacteriaceae family which produce the desired L-amino acid and in which at least one or more of the genes of the cysteine biosynthesis pathway chosen from the group consisting of cysG, cysB, cysZ, cysK, cysM, cysA, cysW, cysU, cysP, cysD, cysN, cysC, cysJ, cysI, cysH, cysE and sbp, or nucleotide sequences which code for these, is (are) enhanced, in particular over-expressed,
 - 10 b) concentration of the desired L-amino acid in the medium or in the cells of the microorganisms, and
 - 15 c) isolation of the desired L-amino acid, constituents of the fermentation broth and/or the biomass in its entirety or portions (> 0 to 100%) thereof optionally remaining in the product.
- 20 2. A process as claimed in claim 1, wherein microorganisms in which further genes of the biosynthesis pathway of the desired L-amino acid are additionally enhanced are employed.
- 25 3. A process as claimed in claim 1, wherein microorganisms in which the metabolic pathways which reduce the formation of the desired L-amino acid are at least partly eliminated are employed.
- 30 4. A process as claimed in claim 1, wherein the expression of the polynucleotide (s) which code(s) for one or more of the genes of cysteine biosynthesis chosen from the group consisting of cysG, cysB, cysZ, cysK, cysM, cysA,

cysW, cysU, cysP, cysD, cysN, cysC, cysJ, cysI, cysH, cysE and sbp is increased.

5. A process as claimed in claim 1, wherein the regulatory and/or catalytic properties of the polypeptides (proteins) for which the polynucleotides cysG, cysB, cysZ, cysK, cysM, cysA, cysW, cysU, cysP, cysD, cysN, cysC, cysJ, cysI, cysH, cysE and sbp code are improved or increased.
10. A process as claimed in claim 1, wherein, for the preparation of L-amino acids, microorganisms of the Enterobacteriaceae family in which in addition at the same time one or more of the genes chosen from the group consisting of:
 15. 6.1 the thrABC operon which codes for aspartate kinase, homoserine dehydrogenase, homoserine kinase and threonine synthase,
 - 6.2 the pyc gene which codes for pyruvate carboxylase,
 20. 6.3 the pps gene which codes for phosphoenol pyruvate synthase,
 - 6.4 the ppc gene which codes for phosphoenol pyruvate carboxylase,
 - 6.5 the pntA and pntB genes which code for transhydrogenase,
 25. 6.6 the rhtB gene which imparts homoserine resistance,
 - 6.7 the mqo gene which codes for malate:quinone oxidoreductase,
 - 6.8 the rhtC gene which imparts threonine resistance.

6.9 the thrE gene which codes for the threonine export protein

6.10 the gdhA gene which codes for glutamate dehydrogenase

5 6.11 the hns gene which codes for the DNA-binding protein HLP-II,

6.12 the pgm gene which codes for phosphoglucomutase,

10 6.13 the fba gene which codes for fructose biphosphate aldolase,

6.14 the ptsH gene which codes for the phosphohistidine protein hexose phosphotransferase,

15 6.15 the ptsI gene which codes for enzyme I of the phosphotransferase system,

6.16 the crr gene which codes for the glucose-specific IIA component,

6.17 the ptsG gene which codes for the glucose-specific IIBC component,

20 6.18 the lrp gene which codes for the regulator of the leucine regulon,

6.19 the mopB gene which codes for 10 Kd chaperone,

6.20 the ahpC gene which codes for the small sub-unit of alkyl hydroperoxide reductase,

25 6.21 the ahpF gene which codes for the large sub-unit of alkyl hydroperoxide reductase,

is or are enhanced, in particular over-expressed, are provided.

7. A process as claimed in claim 1, wherein, for the preparation of L-amino acids, microorganisms of the Enterobacteriaceae family in which in addition at the same time one or more of the genes chosen from the group consisting of:

5 7.1 the *tdh* gene which codes for threonine dehydrogenase,

10 7.2 the *mdh* gene which codes for malate dehydrogenase,

15 7.3 the gene product of the open reading frame (orf) *yjfA*,

 7.4 the gene product of the open reading frame (orf) *yjfP*,

 7.5 the *pckA* gene which codes for phosphoenol pyruvate carboxykinase,

20 7.6 the *poxB* gene which codes for pyruvate oxidase,

 7.7 the *aceA* gene which codes for isocitrate lyase,

 7.8 the *dgsA* gene which codes for the *DgsA* regulator of the phosphotransferase system,

25 7.9 the *fruR* gene which codes for the fructose repressor,

 7.10 the *rpoS* gene which codes for the sigma³⁸ factor

 is or are attenuated, in particular eliminated or reduced in expression, are fermented.

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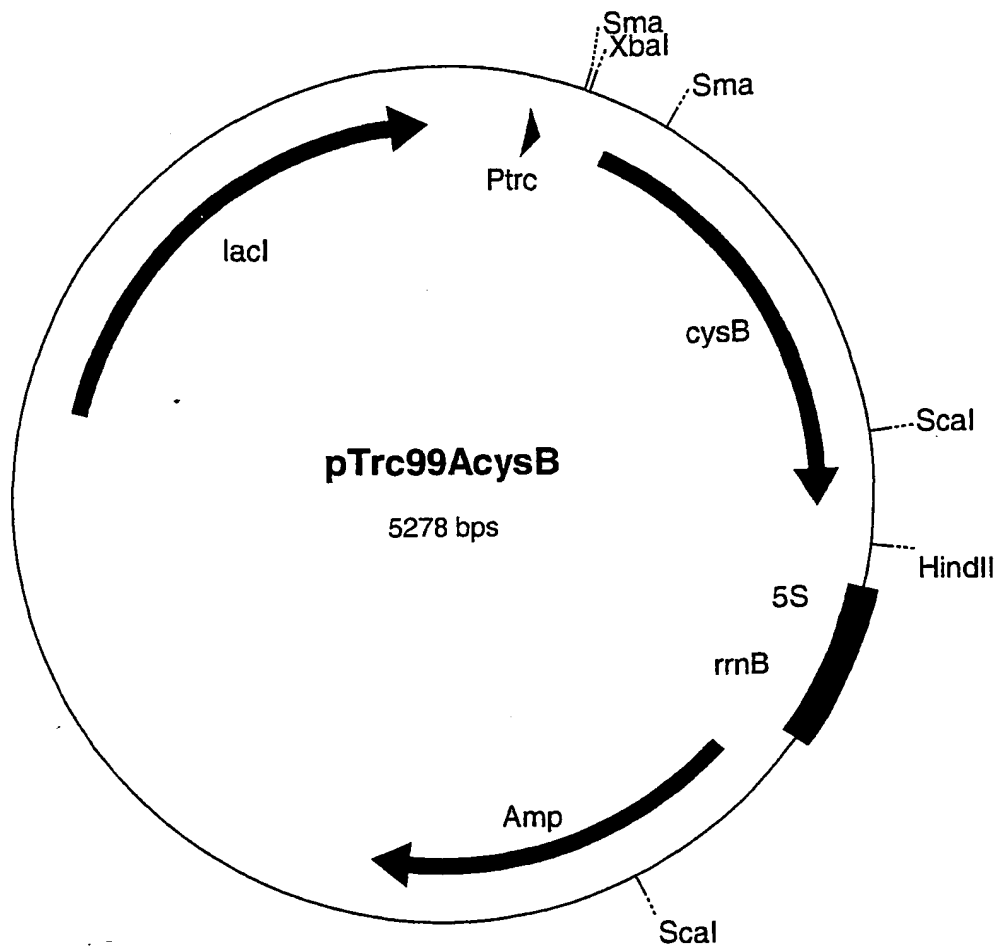


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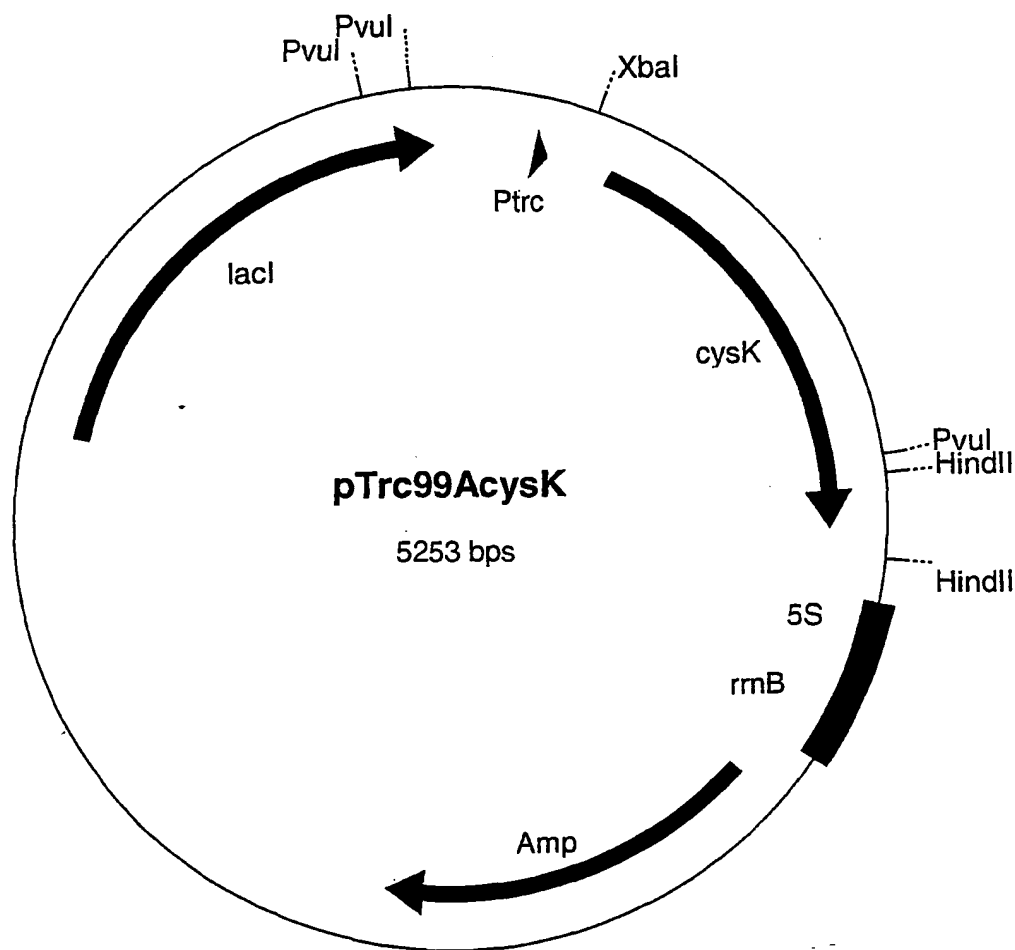


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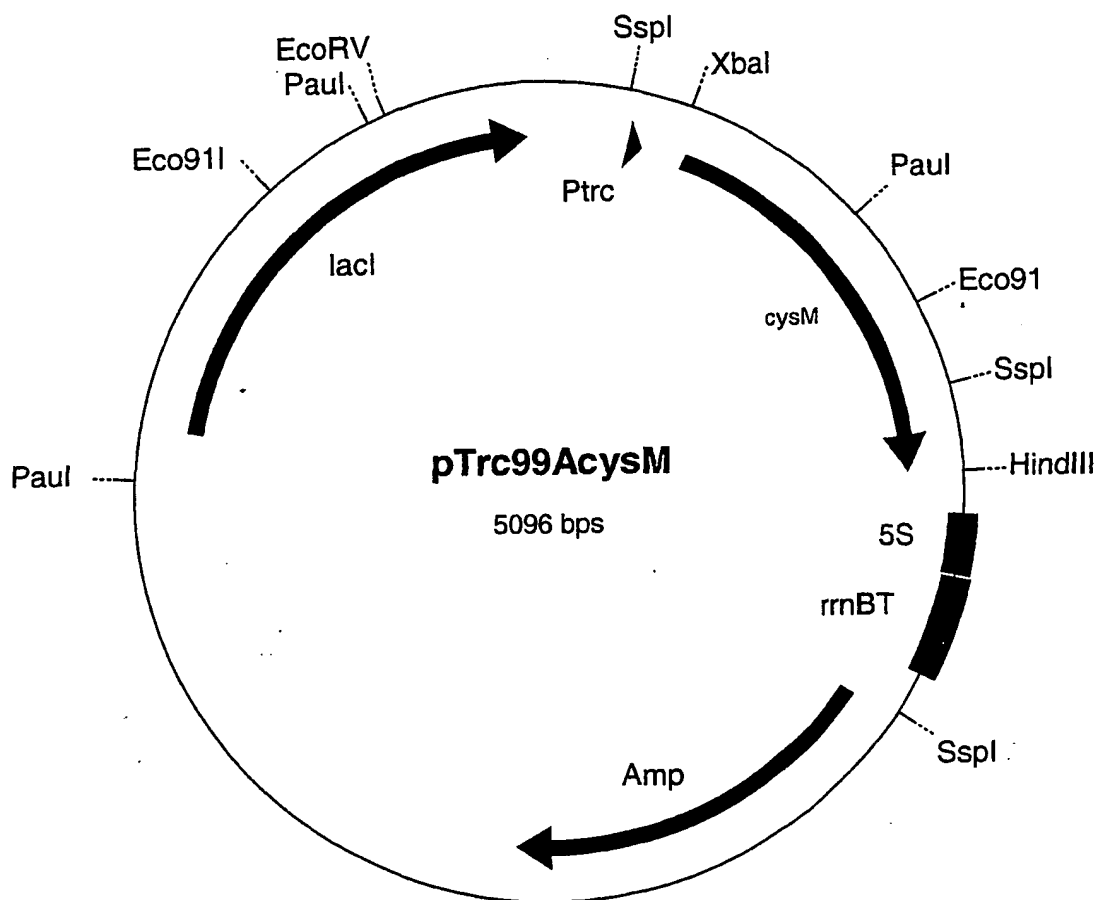


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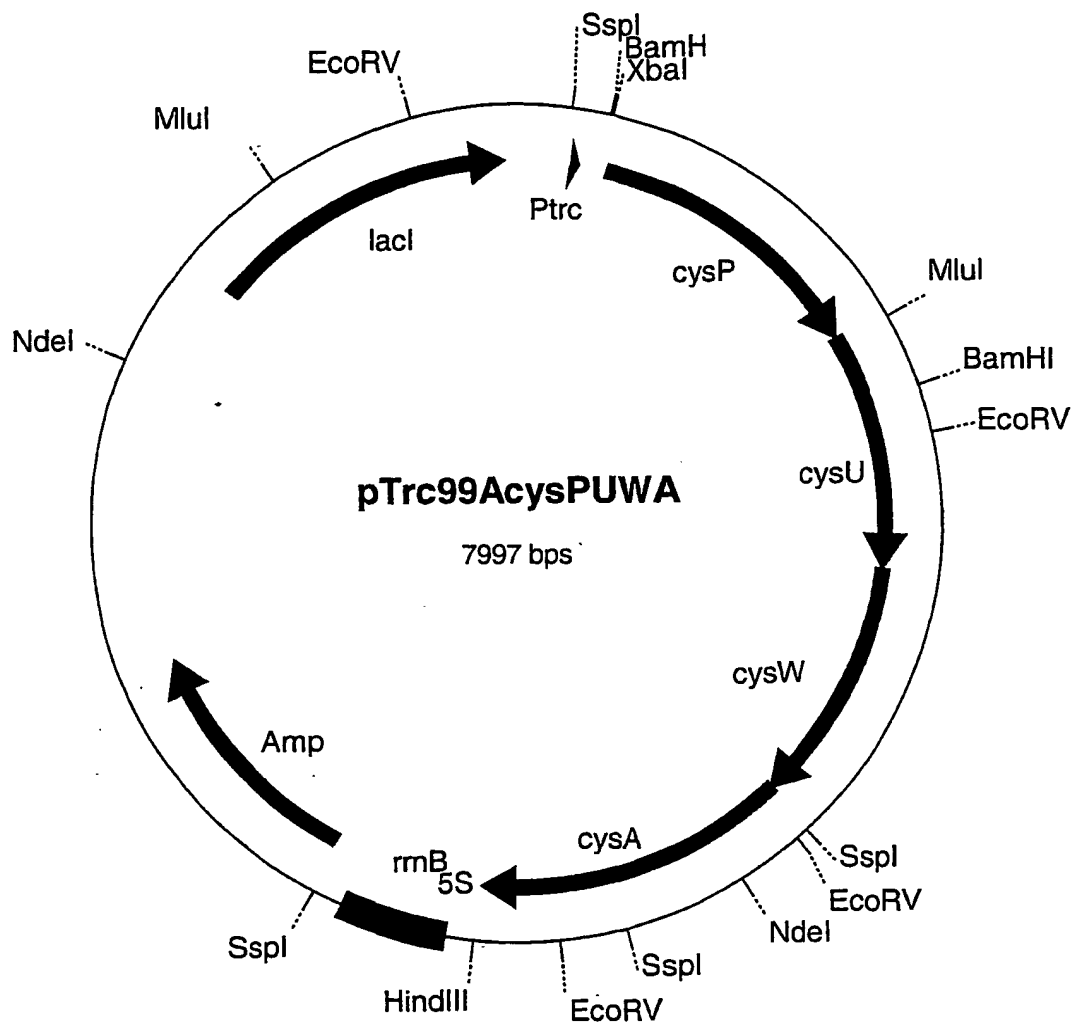


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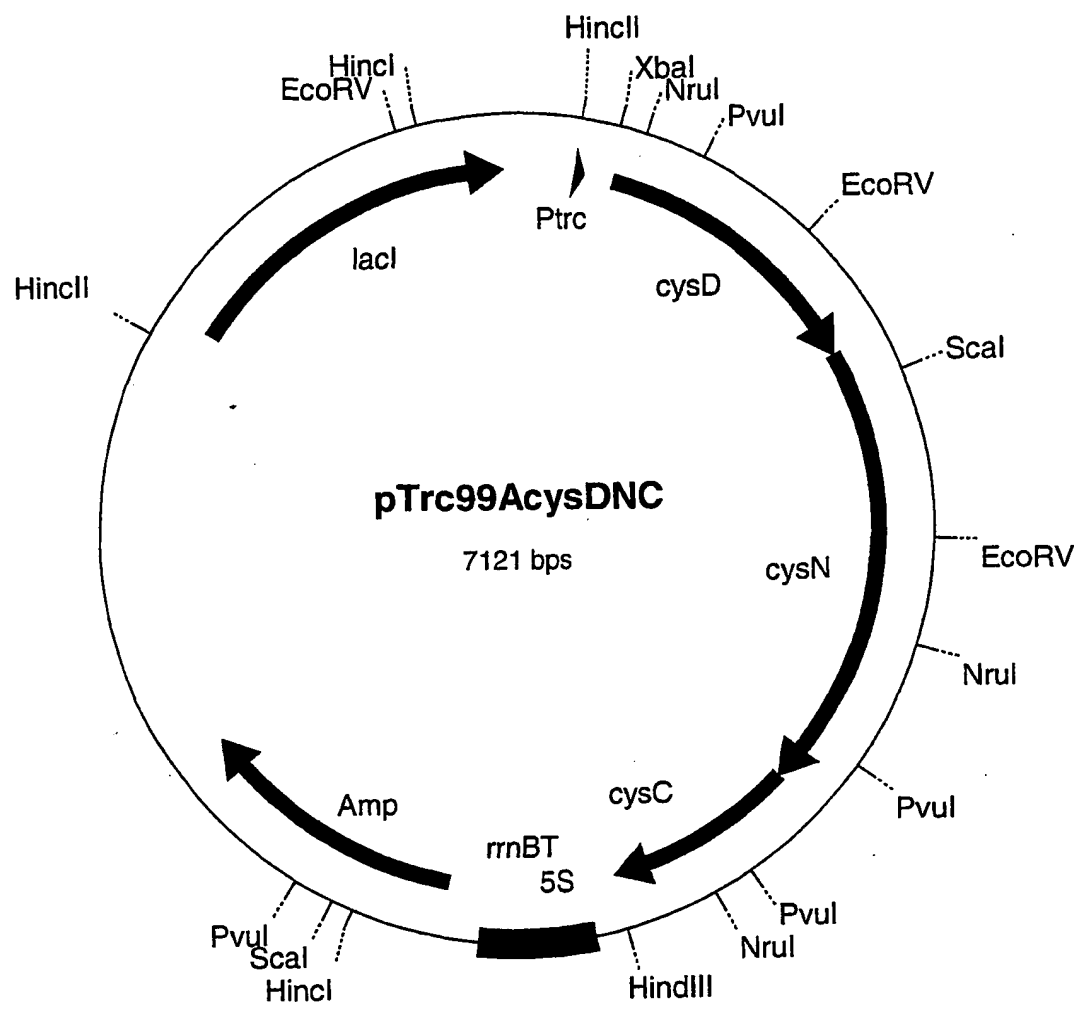


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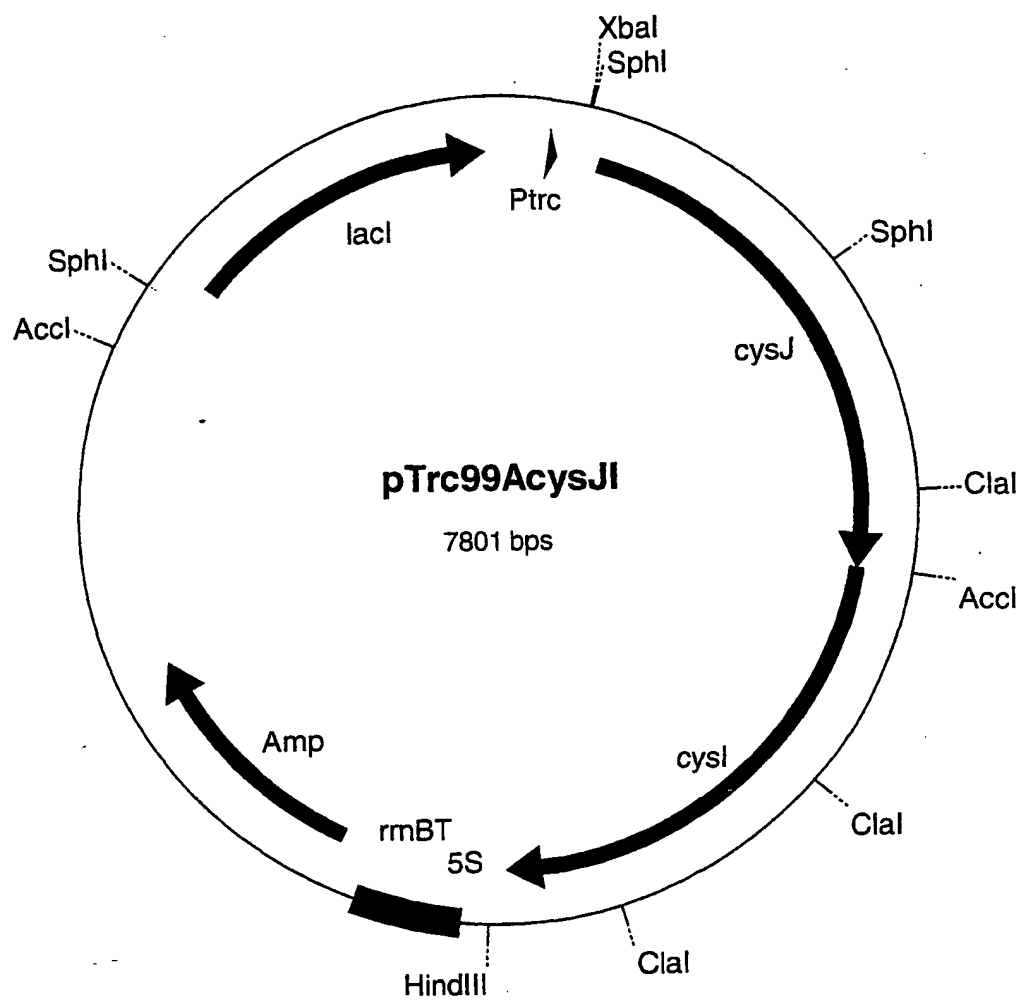
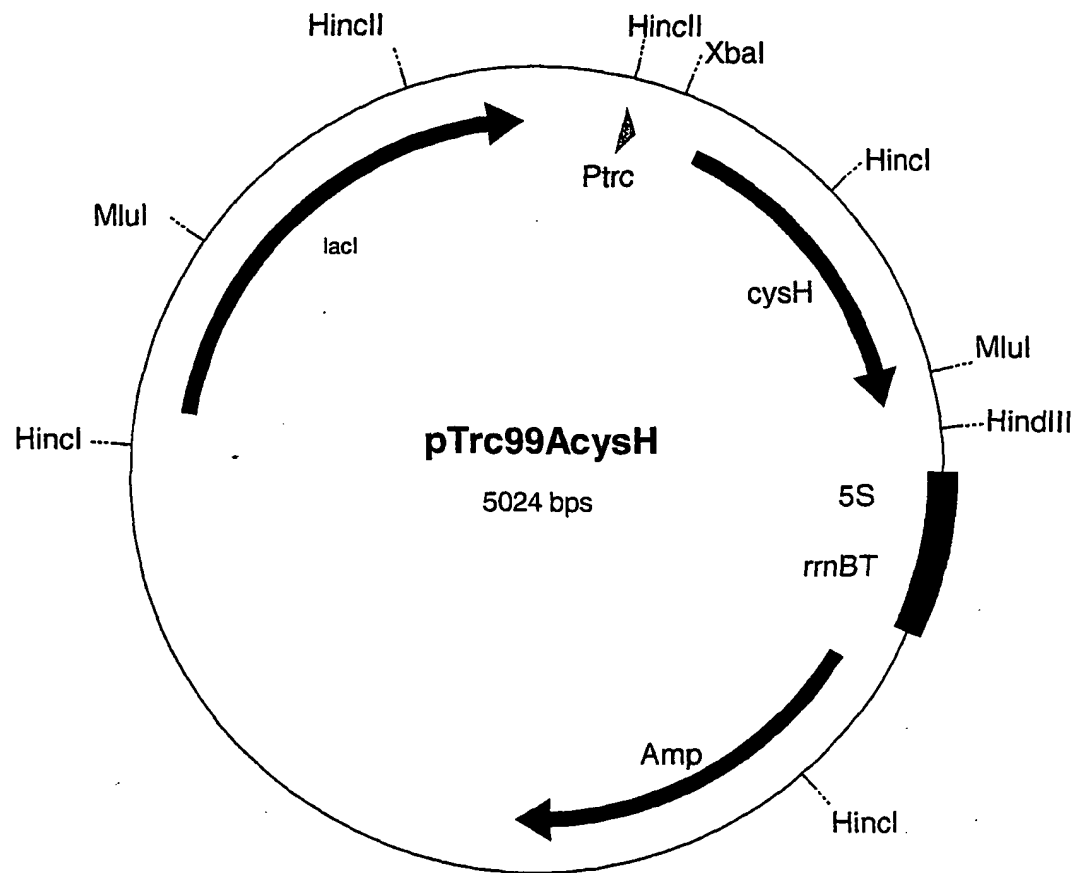


Figure 7:



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LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW,
MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG,
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03/006666 A3

(54) Title: PROCESS FOR THE PREPARATION OF L-AMINO ACIDS USING STRAINS OF THE ENTEROBACTERIACEAE FAMILY

(57) Abstract: The invention relates to a process for the preparation of L-amino acids, in particular L-threonine, in which the following steps are carried out: a) fermentation of microorganisms of the Enterobacteriaceae family which produce the desired L-amino acid, b) separation of the culture medium from the microorganisms, c) addition of a protease to the culture medium, d) hydrolysis of the culture medium, e) separation of the culture medium, f) removal of the culture medium, g) addition of an amino acid oxidase to the culture medium, h) oxidation of the culture medium, i) separation of the culture medium, j) removal of the culture medium, k) addition of an amino acid oxidase to the culture medium, l) oxidation of the culture medium, m) separation of the culture medium, n) removal of the culture medium, o) addition of an amino acid oxidase to the culture medium, p) oxidation of the culture medium, q) separation of the culture medium, r) removal of the culture medium, s) addition of an amino acid oxidase to the culture medium, t) oxidation of the culture medium, u) separation of the culture medium, v) removal of the culture medium, w) addition of an amino acid oxidase to the culture medium, x) oxidation of the culture medium, y) separation of the culture medium, z) removal of the culture medium.

INTERNATIONAL SEARCH REPORT

International Application No PCT/EP 02/06187

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C12P13/06 C12P13/08 C12P13/10 C12P13/12 C12P13/14 C12P13/20 C12P13/22 C12P13/24											
According to International Patent Classification (IPC) or to both national classification and IPC											
B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC 7 C12P											
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched											
Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, WPI Data, PAJ, BIOSIS											
C. DOCUMENTS CONSIDERED TO BE RELEVANT <table border="1" style="width: 100%; border-collapse: collapse;"> <thead> <tr> <th style="text-align: left; padding: 2px;">Category °</th> <th style="text-align: left; padding: 2px;">Citation of document, with indication, where appropriate, of the relevant passages</th> <th style="text-align: left; padding: 2px;">Relevant to claim No.</th> </tr> </thead> <tbody> <tr> <td style="padding: 2px;">A</td> <td style="padding: 2px;"> PEAKMAN T ET AL: "NUCLEOTIDE SEQUENCE, ORGANISATION AND STRUCTURAL ANALYSIS OF THE PRODUCTS OF GENES IN THE NLRB - CYSG REGION OF THE ESCHERICHIA COLI K-12 CHROMOSOME" EUROPEAN JOURNAL OF BIOCHEMISTRY, BERLIN, DE, vol. 191, no. 2, July 1990 (1990-07), pages 315-323, XP001030905 ISSN: 0014-2956 cited in the application the whole document --- -/-/ </td> <td style="padding: 2px;">1-7</td> </tr> </tbody> </table>						Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	A	PEAKMAN T ET AL: "NUCLEOTIDE SEQUENCE, ORGANISATION AND STRUCTURAL ANALYSIS OF THE PRODUCTS OF GENES IN THE NLRB - CYSG REGION OF THE ESCHERICHIA COLI K-12 CHROMOSOME" EUROPEAN JOURNAL OF BIOCHEMISTRY, BERLIN, DE, vol. 191, no. 2, July 1990 (1990-07), pages 315-323, XP001030905 ISSN: 0014-2956 cited in the application the whole document --- -/-/	1-7
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<input checked="" type="checkbox"/> Further documents are listed in the continuation of box C.			<input checked="" type="checkbox"/> Patent family members are listed in annex.								
° Special categories of cited documents : "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed											
"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family											
Date of the actual completion of the international search 16 March 2003			Date of mailing of the international search report 03.06.2003								

INTERNATIONAL SEARCH REPORT

International Application No

PCT/EP 02/06187

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	WARREN M J ET AL: "Gene dissection demonstrates that the <i>Escherichia coli</i> cysG gene encodes a multifunctional protein." THE BIOCHEMICAL JOURNAL. ENGLAND, vol. 302 (Pt 3), 15 September 1994 (1994-09-15), pages 837-844, XP008014987 ISSN: 0264-6021 the whole document ---	1-7
A	WOODCOCK S C ET AL: "Effect of mutations in the transmethylase and dehydrogenase/chelatase domains of sirohaem synthase (CysG) on sirohaem and cobalamin biosynthesis." THE BIOCHEMICAL JOURNAL. ENGLAND, vol. 330 (Pt 1), 15 February 1998 (1998-02-15), pages 121-129, XP002234624 ISSN: 0264-6021 the whole document ---	1-7
A	DE 199 49 579 C (CONSORTIUM ELEKTROCHEM IND) 16 November 2000 (2000-11-16) the whole document ---	1-7
A	DE 195 39 952 A (CONSORTIUM ELEKTROCHEM IND) 30 April 1997 (1997-04-30) the whole document ---	1-7
A	OKAMOTO K ET AL: "HYPERPRODUCTION OF L-THREONINE BY AN <i>ESCHERICHIA COLI</i> MUTANT WITH IMPAIRED L-THREONINE UPTAKE" BIOSCIENCE BIOTECHNOLOGY BIOCHEMISTRY, JAPAN SOC. FOR BIOSCIENCE, BIOTECHNOLOGY AND AGROCHEM. TOKYO, JP, vol. 61, no. 11, November 1997 (1997-11), pages 1877-1882, XP001018682 ISSN: 0916-8451 the whole document -----	1-7

INTERNATIONAL SEARCH REPORT

International application No.
PCT/EP 02/06187

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:

2. Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:

3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

see additional sheet

1. As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.

2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.

3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:

4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

1-7 partial

Remarks on Fees

The additional search fees were deposited in time and in full.

No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1-7 partial

Processes for the preparation of L-amino acids, which comprises fermentation of microorganism of the Enterobacteriaceae family which produce the desired L-amino acid in which the cysG gene, or the polynucleotide which code for this, is enhanced, or wherein the regulatory and/or catalytic properties of said gene are improved. Said process wherein at least one other gene selected from the group cysB, cysZ(K), cysM, cysA, cysW, cysU, cysP(J), cysD, cysN, cysC, cysI, cysH, cysE, or sbp is enhanced. A process wherein further genes of the biosynthesis pathway for the desired amino acid are additionally enhanced. Process wherein microorganisms are employed in which, in addition to the enhanced cysG gene, the metabolic pathways which reduce the formation of the desired amino acid are at least partly eliminated. Process wherein, in addition to the enhanced cysG gene, one or more additional amplified genes are selected from the group: thrABC, pyc, pps, ppc, pntA, pntB, rhtB, mqo, rhtC, thrE, gdhA, hns,pgm, fba, ptsH, ptsI, crr, ptsG, lrp, mopB, ahpC and/or ahpF. Process wherein in addition to the enhanced cysG gene, one or more attenuated genes are selected from the group: tdh, mdh, yjfa, yjfP, pckA, poxB, aceA, dgsA, fruR, and/or rpoS.

2. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysB gene.

3. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysZ (identical to the cysK) gene.

4. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysM gene.

5. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysA gene.

6. Claims: 1-7 partial

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

See subject 1, replacing in the text the cysG for the cysW gene.

7. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysU gene.

8. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysP (identical to the cysJ) gene.

9. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysD gene.

10. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysN gene.

11. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysC gene.

12. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysI gene.

13. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysH gene.

14. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysE gene.

15. Claims: 1-7 partial

See subject 1, replacing in the text the cysG for the cysF gene.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

gene.